AMENDMENTS TO THE CLAIMS

1-48 (Canceled).

- 49 (Currently Amended). A method for treating on-demand at least one symptom of gastro-esophageal reflux disease (GERD) in a human comprising
- (i) identifying a proton pump inhibitor or a salt thereof (PPI) selected from a group consisting essentially of acid-activated agents that inhibit the gastric H+,K+-ATPase enzyme lansoprazole, omeprazole, pantoprazole, rabeprazole, pariprazole, leminoprazole, and their pharmaceutically acceptable salts, isomers including enantiomers, and pharmaceutically acceptable salts of said isomers,
- (ii) identifying an H2 receptor antagonist or a salt therefore (H2RA) selected from a group consisting essentially of agents that inhibit action of histamine on H2 receptors on parietal cell surfaces cimetidine, ranitidine, nizatidine and famotidine, and their pharmaceutically acceptable salts, isomers, and pharmaceutically acceptable salts of said isomers,
 - (iii) adopting an oral dose regime comprising:
 - (a) selecting an oral dosage form for the H2RA for release of H2RA in a gastrointestinal tract;
 - (b) selecting an oral dosage form for the PPI for release of PPI in the gastro-intestinal tract and that, when orally administered to the gastro-intestinal tract concomitantly concurrently with the H2RA, delays and/or extends the release of the PPI relative to the release of the H2RA;
 - (c) orally administering concomitantly the selected oral dosage forms of the H2RA and the PPI to affect a rise in gastrie pH to above about 3 within about 2 hours of administration,
- (iv) on demand, based upon an occurrence of at least one symptom of GERD, orally administering the selected oral dosage forms of the PPI and the H2RA eoneomitantly concurrently according to the dose regime to affect a rise in gastric pH to above about 3 within about 2 hours of administration, thereby treating at least one symptom of GERD promptly, and
- (v) repeating (iv) on demand, based upon a subsequent occurrence of at least one symptom of GERD, if necessary over a prolonged period until 6 hours from the administration of the last dose.

wherein the at least one symptom of GERD is selected from a group consisting of heartburn, sour stomach, and upper abdominal pain.

50-86 (Canceled).

- 87 (Currently Amended). A method as claimed in claim 49, wherein (iii)(a) and (iii)(b) comprise selecting separate oral dosage forms for the H2RA and the PPI, and wherein iii(c) and (iv) comprises orally administering the separately oral dosage forms concomitantly on demand concurrently.
- 88 (Currently Amended). A method as claimed in claim 87, wherein the separate oral dosage form for the PPI comprises is a tablet or capsule within which the PPI is presented as a plurality of small dosage units comprising pellets, granules or beads distributed within the tablet or capsule.
- 89 (Currently Amended). A method as claimed in claim 49, wherein (iii)(a) and (iii)(b) comprise combining the oral dosage forms for the H2RA and the PPI into a single oral dosage form, and wherein (iii)(c) and (iv) comprises orally administering the single oral dosage form on demand.
- 90 (Currently Amended). A method as claimed in claim 88 89, wherein the single oral dosage form comprises is a tablet or capsule within which the PPI is presented as a plurality of small dosage units comprising pellets, granules or beads distributed within the tablet or capsule.
- 91 (Previously Submitted). A method as claimed in claim 90, wherein the tablet or capsule further comprises a pharmaceutically acceptable excipient.
- 92 (Previously Submitted). A method as claimed in claim 91, wherein the pharmaceutically acceptable excipient comprises a disintegrant.
 - 93 (Cancelled).
- 94 (Currently Amended). A method as claimed in claim 89, wherein the single oral dosage form includes a core comprising the PPI and a membrane including an excipient applied onto the core that delays and/or extends the release of the PPI relative to the release of the H2RA.
- 95 (Previously Submitted). A method as claimed in claim 94, wherein the single oral dosage form includes an alkaline-reacting substance admixed with the PPI.
- 96 (Currently Amended). A method as claimed in claim 94, wherein the single oral dosage form comprises is presented as a tablet or capsule within which the core and the membrane

are presented as a plurality of small dosage units comprising pellets, granules or beads distributed within the tablet or capsule.

- 97 (Currently Amended). A method as claimed in claim 94, wherein the single oral dosage form comprises is presented in two halves, one of which comprises one or more of the cores and membranes, and the other half comprises the H2RA.
- 98 (Previously Submitted). A method as claimed in claim 97, wherein the H2RA half includes a pharmaceutically acceptable excipient.
- 99 (Previously Submitted). A method as claimed in claim 98, wherein the pharmaceutically acceptable excipient comprises a disintegrant.
- 100 (Previously Submitted)). A method as claimed in claim 97 or 98, wherein the H2RA forms an outer layer applied onto the membrane of the core.
- 101 (Previously Submitted). A method as claimed in claim 94, wherein the H2RA forms an outer layer applied onto the membrane of the core, and wherein the single dosage form includes an alkaline-reacting substance admixed with the PPI.
- 102 (Previously Submitted)). A method as claimed in claim 94, wherein the single dosage form includes an enteric coating layer applied onto the membrane.
- 103 (Previously Submitted). A method as claimed in claim 102, wherein the single dosage form includes a layer separating the enteric coating from the membrane.
- 104 (Previously Submitted). A method as claimed in claim 102 or 103, wherein the H2RA forms an outer layer applied onto the membrane of the core.
- 105 (Currently Amended). A method as claimed in claim 89, wherein the PPI of the single oral dosage form includes is presented as a matrix comprising the PPI and an excipient incorporated with the PPI to delay and/or extend the release of the PPI relative to the release of the H2RA.
- 106 (Previously Submitted). A method as claimed in claim 105, wherein the single oral dosage form includes an alkaline-reacting substance admixed with the PPI.
- 107 (Currently Amended). A method as claimed in claim 105, wherein the single oral dosage form comprises is a tablet or capsule within which the matrix is presented as a plurality of small dosage units comprising pellets, granules or beads distributed within the tablet or capsule.

- 108 (Previously Submitted). A method as claimed in claim 105 or 107, wherein the H2RA forms an outer layer applied onto the matrix.
- 109 (Previously Submitted). A method as claimed in claim 105, wherein the H2RA forms an outer layer applied onto the matrix, and wherein the single oral dosage form includes an alkaline-reacting substance admixed with the PPI.
- 110 (Currently Amended). A method as claimed in claim 105, wherein the single oral dosage form comprises is presented in two halves, one of which comprises the matrix, and the other half comprises the H2RA.
- 111 (Previously Submitted). A method as claimed in claim 110, wherein the H2RA half includes a pharmaceutically acceptable excipient.
- 112 (Previously Submitted). A method as claimed in claim 111, wherein the pharmaceutically acceptable excipient comprises a disintegrant.
- 113 (Previously Submitted). A method as claimed in claim 105, wherein the single dosage form includes an enteric coating layer applied onto the matrix.
- 114 (Previously Submitted). A method as claimed in claim 113, wherein the single dosage form includes a layer separating the enteric coating from the matrix.
- 115 (Previously Submitted). A method as claimed in claim 113 or 114, wherein the H2RA forms an outer layer applied onto the matrix.
- 116 (Currently Amended). A method as claimed in claim 49, wherein at least one of the selected oral dosage forms further comprises an antacid agent and/or or an alginate.
- 117 (Currently Amended). A method as claimed in claim 116, wherein the antacid agent comprises at least one of aluminum hydroxide, calcium carbonate, magnesium carbonate, basic magnesium carbonate, magnesium hydroxide, magnesium oxide and/or or sodium hydrogen carbonate.